### 1) Basics

- 1.1) Introduction to standard MPT models
- 1.2) Examples
- 1.3) Model development
- 1.4) Formal model structure
- 1.5) Identifiability
- 1.6) Parameter estimation
- 1.7) Model assessment
- 1.8) Selected literature

### 1.4) Formal model structure

- Multinomial models
- Parameterized multinomial models
- Multinomial processing tree (MPT) models
- Processing-tree representation of MPT models

#### Multinomial Models

- *a)* One condition (simple multinomial model)
- One variable with J categories and sample frequencies  $n_1, n_2, ..., n_j, ..., n_J$ .
- $\pi = (p_1, p_2, ..., p_J)$  is the vector of category probabilities.
- Given independent sampling, the sample frequencies follow a multinomial distribution:

$$p_{N,\pi}(n_1,n_2,...,n_J) = \frac{N}{n_1! n_2! ... n_J!} p_1^{n_1} p_2^{n_2} ... p_J^{n_J}$$

#### b) Several conditions (joint multinomial model)

- In each condition k (k = 1, ..., K), one categorical variable with J(k) categories is observed.
- For each of the *K* conditions a simple multinomial model holds.
- Given independence between conditions, the overall probability of the sample frequencies across conditions is

$$p = \prod_{k=1}^{K} p_{N(k),\pi(k)}(n_{1(k)}, n_{2(k)}, ..., n_{J(k)})$$

#### Parameterized Multinomial Models

- The category probabilities  $p_1, p_2$  etc. are rewritten as functions of "latent parameters"  $\theta_1, \theta_2, ..., \theta_S$
- Thus, in case of a simple multinomial model we have

$$- p_1 = f_1(\theta_1, \theta_2, ..., \theta_S)$$

$$- p_2 = f_2(\theta_1, \theta_2, ..., \theta_S)$$

$$- ....$$

$$- p_J = f_J(\theta_1, \theta_2, ..., \theta_S)$$

- These equations are called model equations.
- The set of possibles values of S latent parameters is called "parameter space"  $\Omega$  of the model.

#### Multinomial Processing Tree (MPT) Models

- MPT models form a subclass of parameterized multinomial models.
- Additional assumptions:
- 1) Each  $\theta_s$  is in [0, 1]

#### Multinomial Processing Tree (MPT) Models

- MPT models form a subclass of parameterized multinomial models.
- Additional assumptions:
- 1) Each  $\theta_s$  is in [0, 1]
- 2) Structure of the model equations (Hu & Batchelder, 1994):

$$p_{j} = \sum_{i=1}^{I(j)} c_{ij} \prod_{s=1}^{S} \theta_{s}^{a_{ijs}} \cdot (1 - \theta_{s})^{b_{ijs}}, \qquad \sum_{j=1}^{J} p_{j} = 1, \qquad \theta_{s} \in [0, 1]$$

where s: Parameter index

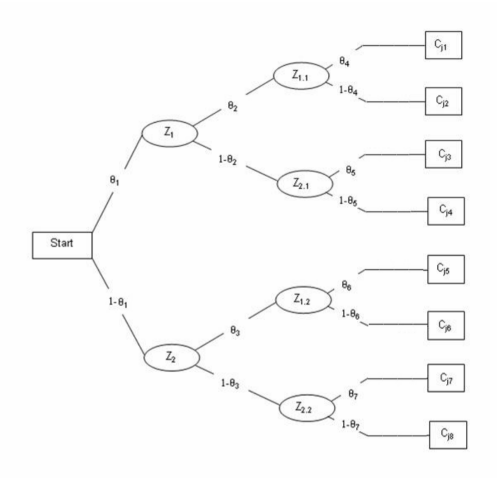
*j* : Category index

*i* : Branch index

 $c_{ii}$ : positive real number

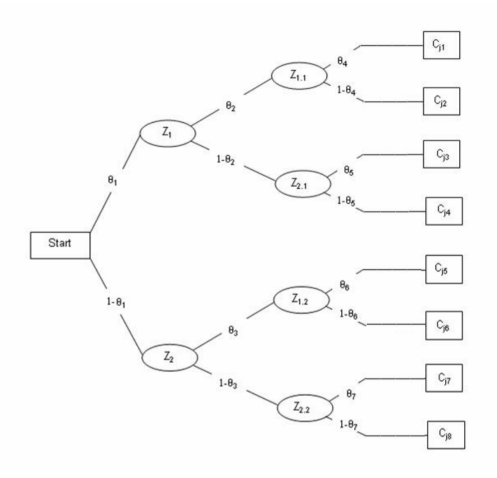
 $a_{iis}$ ,  $b_{iis}$ : nonnegative integer number (often 0 or 1)

## Binary probabilistic event trees can always be translated in MPT-model equations:



#### ... as an aside ...

- Although any binary processing tree diagram uniquely determines a system of MPT model equations ...
- ... it is not true that any system of MPT model equations uniquely determines a specific processing tree diagram.
- Counter examples:
  - Level switching in independence models
  - Category switching given identical branch probabilities



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Foundations II -- MPT Modeling (Erdfelder & Heck)

# Another counter example is this MPT model:

$$p_1(\theta) = \theta^3$$

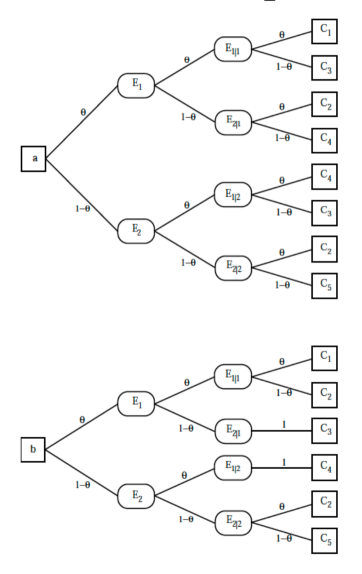
$$p_2(\theta) = \theta \cdot (1 - \theta)$$

$$p_3(\theta) = \theta \cdot (1 - \theta)$$

$$p_4(\theta) = \theta \cdot (1 - \theta)$$

$$p_5(\theta) = (1 - \theta)^3$$

## Two different processing trees compatible with the model on the previous slide:



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## 1.5) Identifiability

Any MPT model equation system

$$p_{j} = \sum_{i=1}^{I(j)} c_{ij} \prod_{s=1}^{S} \theta_{s}^{a_{ijs}} \cdot (1 - \theta_{s})^{b_{ijs}}, \qquad \sum_{j=1}^{J} p_{j} = 1, \qquad \theta_{s} \in [0, 1]$$

where s: Parameter index

*j* : Category index

*i*: Branch index

 $c_{ii}$ : positive real number

 $a_{ijs}$ ,  $b_{ijs}$ : nonnegative integer number (often 0 or 1)

defines a mapping  $f: \Omega \rightarrow P$ 

## Parameter Space and Data Space

- $\Omega$  is called "Parameter Space":
  - = Set of all possible parameter vectors

- P is called "Data Space" (more precisely: space of category probabilities)
  - = Set of all possible category probability vectors

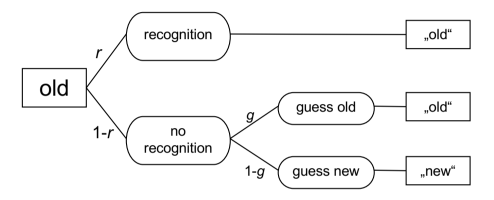
Definition (global identifiability):

A MPT model is globally identified if f is one-to-one.

Definition (local identifiability):

A MPT model is locally identified if f is one-to-one in the neighborhood of  $\theta_0$  in  $\Omega$ .

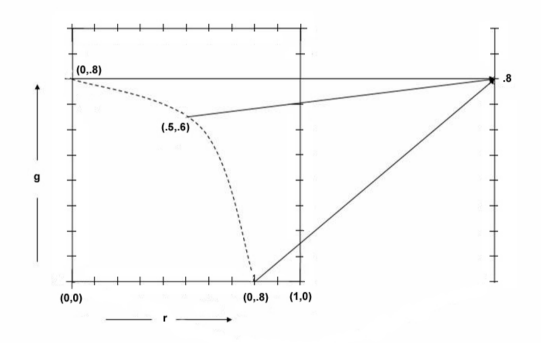
## One-High-Threshold recognition model (Blackwell, 1953)



r: probability of recognition

g: probability of guessing old given recognition

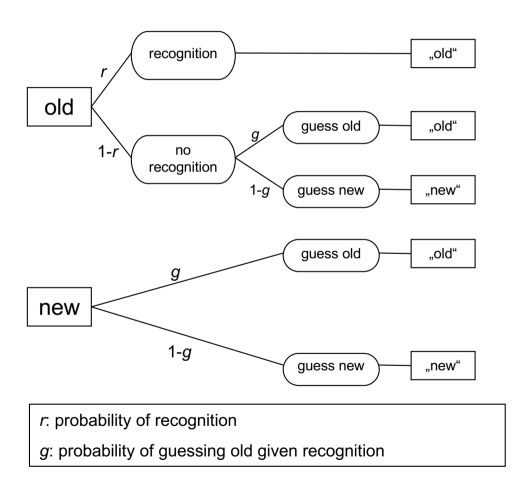
#### Example 1 (nonidentifiability)



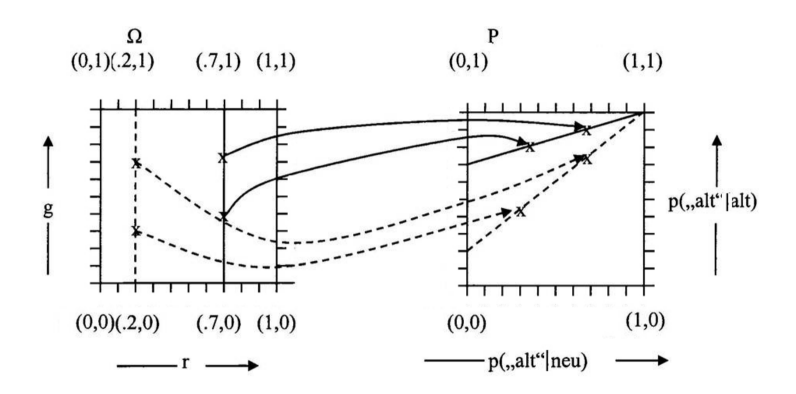
- One-high-Threshold-Modell limited to old items:
  - p("alt" | old item) =  $r + (1-r) \cdot g$
- Model is not identified.

## One-High-Threshold recognition model

(Blackwell, 1953)



#### Example 2 (identifiability)



- One-high-Threshold-Model (old and new items)
  - p("alt" | old item) =  $r + (1-r) \cdot g$
  - $p(,,alt" \mid new item) = g$
- Model ist globally identified.

#### Two important theorems:

• "Observable branches":

A model is always globally identified if each of its branches terminates in a unique empirical category (Hu & Batchelder, 1994).

#### Two important theorems:

• "Observable branches":

A model is always globally identified if each of its branches results in a new empirical category (Hu & Batchelder, 1994).

• "No more parameters than degrees of freedom in the data":

A necessary but not sufficient condition of identifiability is

$$S \leq \sum_{k=1}^{K} (J(k) - 1).$$

#### Jacobian Matrix

- Jacobian: Matrix of the 1st partial derivates of the model equations with respect to parameters  $\theta_s$ , s = 1, ... S.
- r: maximum rank of the Jacobian across  $\Omega$
- If r < S, then the model is neither locally nor globally identified.
- If r = S, then the model is locally identified (but not necessarily globally).

# Remedies for nonidentifiable models

- Parameter constraints
  - Parameter fixations ( $\theta_s = c$ , with c = constant)
  - Equality constraints ( $\theta_s = \theta_{s'}$ )
- Increase the number of empirical categories
  - Additional conditions, no (or few) additional parameters
  - Selective manipulations of parameters

### 1.6) Parameter Estimation

Find a parameter vector  $\mathbf{\theta} = (\theta_1, ..., \theta_s, ..., \theta_s), \mathbf{\theta} \in \Omega$ , such that the distance between the sample frequencies  $n_1, n_2, ..., n_J$  and the expected category frequencies under the model,  $N \cdot p_1(\mathbf{\theta}), N \cdot p_2(\mathbf{\theta}), ..., N \cdot p_J(\mathbf{\theta})$ , becomes a minimum.

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Which distance measure?

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Which distance measure?

The likelihood ratio statistic  $G^2$  and Pearson's  $\chi^2$  are used most often:

$$G^{2}(\theta) = 2\sum_{j=1}^{J} n_{j} \ln \left( \frac{n_{j}}{N \cdot p_{j}(\theta)} \right) \qquad \chi^{2}(\theta) = \sum_{j=1}^{J} \frac{\left[ n_{j} - N \cdot p_{j}(\theta) \right]^{2}}{N \cdot p_{j}(\theta)}$$

Both distance measures are special cases of the Power-Divergence-family ( $PD_{\lambda}$ -Statistics) (Read & Cressie, 1988):

$$s_{\lambda} = \frac{2}{\lambda(\lambda+1)} \sum_{k=1}^{k} \sum_{i=1}^{J(k)} n_{j(k)} \cdot \left[ \left( \frac{n_{j(k)}}{e_{j(k)}} \right)^{\lambda} - 1 \right]$$

Note that:

Pearson-
$$\chi^2$$
 =  $S_{\lambda=1}$   
Likelihood-ratio- $G^2$  =  $\lim_{\lambda \to 0} S_{\lambda}$ .

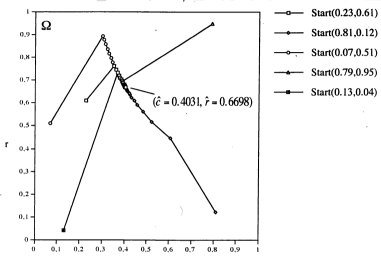
# What is the best estimation method?

- All  $S_{\lambda}$ -estimates have the same asymptotic properties:
  - Consistent, efficient, asympotically unbiased
  - (multivariate) normal sampling distributions
- Choosing  $\lambda$ =0 (i.e., minimizing  $G^2$ ) provides Maximum Likelihood (ML) estimates:
  - Sample frequencies have highest likelihood
  - Standard method in typical applications
- All  $S_{\lambda}$ -estimates are easily implemented using the robust Expectation-Maximization (EM) estimation algorithm.

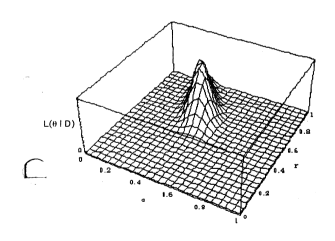
#### Expectation-Maximization-(EM) Algorithm

- 1) Choose a start vector  $\theta_i$ .
- 2) **E**(xpectation)-Step:
  - Estimate the expected frequencies of the branches given  $\theta_i$  and the observed category frequencies  $n_{i(k)}$
- 3) M(aximization)-Step:
  - Let i = i + 1
  - Compute new  $S_{\lambda}$  estimates  $\theta_i$  given the expected frequencies from step 2)
- 4) Convergence ?
  - If  $Abs(\theta_i \theta_{i-1}) > Criterion go back to step 2).$
- 5) Otherwise accept  $\theta_i$  as final parameter estimates  $\hat{\theta}$

## Graphical Illustration of the EM-Algorithm applied to c and r in storage-retrieval model:



EM paths in  $\Omega$  for  $\lambda = 0$ , given five different starting values (simulated data).

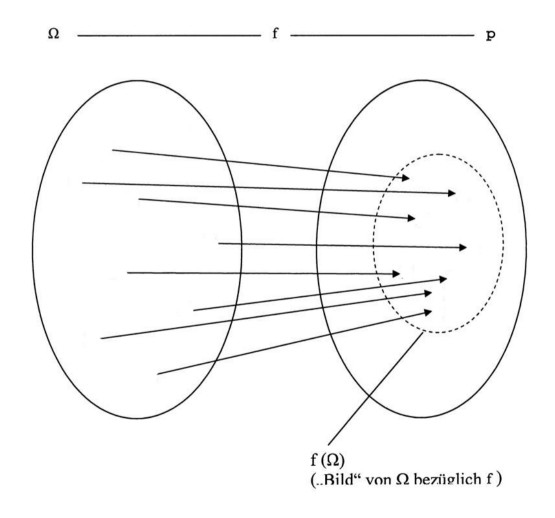


Likelihood (y-axis) as a function of c und r given the same data.

#### Bottom line

- Use  $G^2$ -minimization (= likelihood maximization) as the default estimation method.
- Repeat the estimation process several times using random start values to make sure that you really found a global maximum of the likelihood function.

### 1.7) Model assessment



How to test  $H_0$ :  $\pi \in f(\Omega)$ ?

Options:

Pearson
$$\chi^2 = \sum_{k=1}^K \sum_{j=1}^{J(k)} (n_{j(k)} - e_{j(k)})^2 / e_{j(k)}$$
,

or

Likelihood - ratio - 
$$G^2 = 2 \cdot \sum_{k=1}^K \sum_{j=1}^{J(k)} n_{j(k)} \cdot \ln(n_{j(k)} / e_{j(k)});$$
  
with  $e_{j(k)} = p_{j(k)}(\hat{\boldsymbol{\theta}}) \cdot N(k)$ 

Both statistics are special cases of Read und Cressies (1988) "Power Divergence" – Statistics:

$$S_{\lambda} = \frac{2}{\lambda(\lambda+1)} \sum_{k=1}^{k} \sum_{j=1}^{J(k)} n_{j(k)} * \left[ \left( \frac{n_{j(k)}}{e_{j(k)}} \right)^{\lambda} - 1 \right]$$

Again:

Pearson- $\chi^2 = S_{\lambda=1}$ Likelihood-ratio- $G^2 = \lim_{\lambda \to 0} S_{\lambda}$ .

All  $S_{\lambda}$  – Statistics are asymptotically (i.e., for  $N \rightarrow$  00) chi-square distributed under  $H_0$  (i.e., the model holds) with

df = 
$$\sum_{k=1}^{K} (J(k)-1)-S$$
.

## What is the best goodness-of-fit statistic?

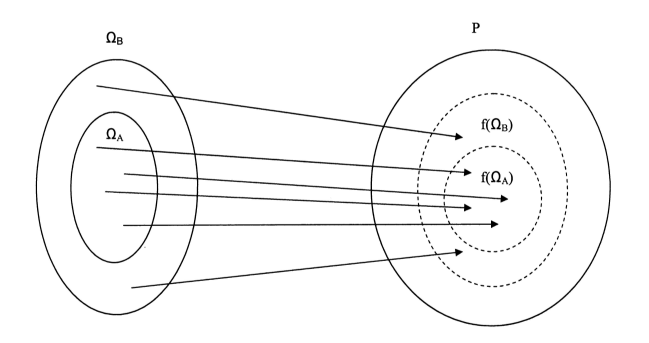
- In case of very small sample sizes,  $S_{\lambda=2/3}$  outperforms other  $S_{\lambda}$ -statistics (such as  $G^2 = S_{\lambda=0}$ ) in terms of accuracy of Chi-square approximation (cf. Read & Cressie, 1988).
- However, very small samples are typically less of a problem in MPT model applications (unless models are tested for single participants).
- Given the fact that  $G^2$  is a by-product of ML-parameter estimation,  $G^2$  can be recommended for moderate to large sample sizes.
- However, you cannot choose  $G^2$  in case of samples with zero cells.
- Remedies: 1) Ignore 0 cells, 2) add constant ε to all counts

## Model comparisons

- Assume that model B  $(M_B)$  holds for your data:  $G^2(M_B)$  is not significant.
- Then often  $M_B$  is compared to an alternative model A  $(M_A)$  that is a special case of  $M_B$ .
- $M_A$  is obtained by applying a parameter restriction to  $M_B$ , e.g., an equality constraint or a parameter fixation.
- How to decide whether  $M_A$  fits the data worse than  $M_B$ ?

Model  $M_A$  as a special case of  $M_B$ :

Illustration of model-specific mapping of  $\Omega_A$  and  $\Omega_B$  in P.



This is called a "hierarchical model family".

# Model comparisons in hierarchical model families

• If both M<sub>A</sub> and M<sub>B</sub> hold then

G<sup>2</sup><sub>A</sub> is chi-square distributed with df<sub>A</sub>

G<sup>2</sup><sub>B</sub> is chi-square distributed with df<sub>B</sub>

 $\Delta G^2_{A-B}$ : =  $G^2_A - G^2_B$  is chi-square distributed with  $df_{A-B} = df_A - df_B$ .

• In other words,  $G^2$  is additive (irrespective of N):

$$G_A^2 = G_B^2 + \Delta G_{A-B}^2$$

- This additivity property does not hold for other  $PD_{\lambda}$ -stats.
- Using  $\Delta G^2_{A-B}$ , it is easy to compare different models in a hierarchical model family using chi-square tests
- This is perhaps the strongest argument for relying on  $G^2$  for purposes of model fitting and testing.

# Model comparisons in nonhierarchical model families

- Unfortunately,  $\Delta G^2_{A-B}$ : =  $G^2_A G^2_B$  cannot be used in nonhierarchical model families.
- How to proceed then?

# Information-theoretic measures of goodness-of-fit

- Akaike Information Criterion (AIC):
  - $AIC(\mathbf{M}_0) = -2 \cdot ln(\mathbf{L}(\mathbf{\theta}; \mathbf{y})) + 2 \cdot S$
  - $-\Delta AIC(M_0) = AIC(M_0)-AIC(sat.) = G^2(M_0) 2 \cdot df(M_0)$
- Bayesian Information Criterion (BIC):
  - $-\operatorname{BIC}(\mathbf{M}_0) = -2 \cdot \ln(\operatorname{L}(\boldsymbol{\theta}; \mathbf{y})) + S \cdot \ln(N)$
  - $-\Delta BIC(M_0) = BIC(M_0) BIC(sat.) = G^2(M_0) df(M_0) \cdot ln(N)$
- Rule:
  - Choose the model with the smaller AIC / BIC
- Recommendation:
  - Fit is excellent if  $\triangle AIC$  or  $\triangle BIC < 0$ .

### 1.8) Selected literature

- Batchelder, W. H., & Riefer, D. M. (1999). Theoretical and empirical review of multinomial process tree modeling. *Psychonomic Bulletin & Review*, 6, 57-86.
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